

results of BLAST

BLASTX 2.2.9 [May-01-2004]

Reference:

Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402.

RID: 1088028727-32641-46334205553.BLASTQ4

Query=

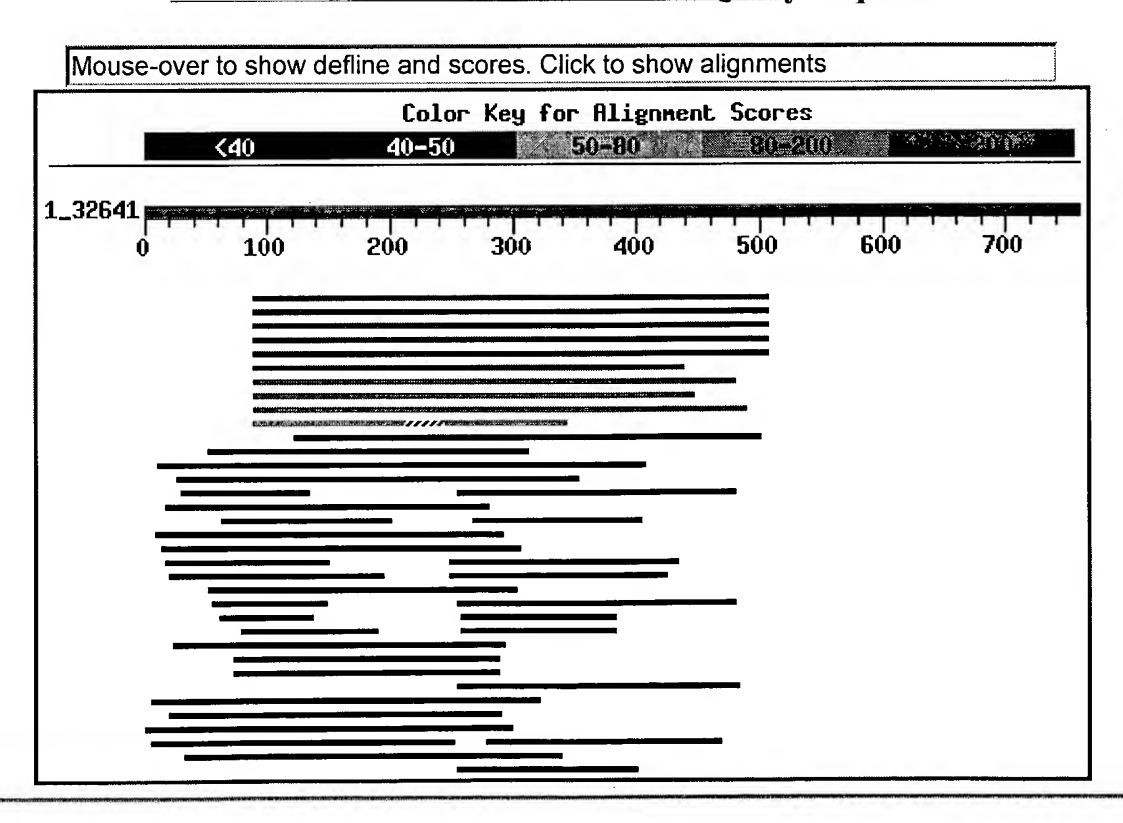
(759 letters)

Database: All non-redundant GenBank CDS translations+PDB+SwissProt+PIR+PRF excluding environmental samples 1,866,121 sequences; 619,474,291 total letters

If you have any problems or questions with the results of this search please refer to the BLAST FAQs

Taxonomy reports

Distribution of 47 Blast Hits on the Query Sequence



Sequences producing significant alignments:	Score (bits)	E Value
gi 34101278 ref NP 898888.1 hypothetical protein MGC10744	280	2e-74 L
gi 34101276 ref NP 115730.2 hypothetical protein MGC10744	272	6e-72 L
gi 21539609 ref NP 080114.1 RIKEN cDNA 1110004B13 [Mus mus	<u> 257</u>	2e-67 L
gi 27672842 ref XP 213332.1 similar to RIKEN cDNA 1110004B	<u> 257</u>	2e-67 L
gi 47123437 gb AAH70231.1 Unknown (protein for MGC:88213)	245	8e-64
gi 21539635 ref NP_082612.1 RIKEN cDNA 1110004B13 [Mus mus	217	2e-55
<u>gi 47086013 ref NP 998376.1 </u> zgc:77926 [Danio rerio] >gi 41 <u>gi 47225380 emb CAG11863.1 </u> unnamed protein product [Tetrao	$\frac{172}{169}$	8e-42 L 4e-41
gi 13751660 gb AAK38512.1 DC20 [Homo sapiens]	<u>119</u>	6e-33
gi 34535000 dbj BAC87177.1 unnamed protein product [Homo s	<u>79</u>	1e-13
gi 7500843 pir T21990 hypothetical protein F39B2.9 - Caeno gi 46362710 ref ZP 00225559.1 hypothetical protein Krad060	$\frac{38}{37}$	0.25 0.33
qi 18025542 gb AAF78882.2 EBNA-3B [cercopithicine herpesvi	37	0.56
qi 41114515 ref XP 371222.1 similar to Hypothetical protei	36	0.73
gi 46321304 ref ZP 00221682.1 COG0477: Permeases of the ma	36	0.95
gi 47210644 emb CAG06315.1 unnamed protein product [Tetrao	35	1.2
gi 34902106 ref NP 912399.1 unknown protein [Oryza sativa	35 35	1.2
gi 7106228 gb AAF36091.1 flagelliform silk protein [Nephil gi 34536289 dbj BAC87601.1 unnamed protein product [Homo s	<u>35</u>	2.1
gi 6686017 sp 062732 SYN1 CANFA Synapsin I >gi 2944066 gb A	35	2.1
gi 14701574 dbj BAB62028.1 elastase precursor [Prevotella	35	2.1
gi 42659679 ref XP_374898.1 similar to pecanex-like 3 [Hom	34	2.8
gi 34365329 emb CAE45990.1 hypothetical protein [Homo sapi	34	2.8 L
gi 46311822 ref ZP 00212424.1 COG0477: Permeases of the ma	34	2.8
gi 13562004 gb AAK30605.1 major ampullate spidroin 2-like	$\frac{34}{34}$	2.8
<pre>gi 28279307 gb AAH46165.1 Unknown (protein for IMAGE:51920 gi 34495966 ref NP 900181.1 probable MFS permease [Chromob</pre>	$\frac{34}{34}$	3.6 3.6
gi 37680831 ref NP 935440.1 hypothetical protein VV2647 [V	34	3.6
gi 27365118 ref NP 760646.1 Unknown [Vibrio vulnificus CMC	34	3.6
gi 24575123 gb AAL06694.1 putative regulatory protein [Str	34	3.6
gi 41057143 ref NP 957857.1 ORF080 virion core protein [Or	34	3.6
gi 47230172 emb CAG10586.1 unnamed protein product [Tetrao	33	4.7
gi 39583515 emb CAE73973.1 Hypothetical protein CBG21601 [33	4.7 6.1
gi 9663052 emb CAC01096.1 110-R orphan receptor [Haemonchu gi 47214200 emb CAG00828.1 unnamed protein product [Tetrao	<u>33</u>	6.1
gi 47217474 emb CAG10243.1 unnamed protein product [Tetrao	<u>33</u> <u>33</u>	6.1
gi 46432590 gb EAK92065.1 hypothetical protein CaO19.6598	33	6.1
gi 15220207 ref NP 172545.1 expressed protein [Arabidopsis	33	8.0
gi 46443901 gb EAL03180.1 hypothetical protein Ca019.4050	33 33 33	8.0
gi 21388708 dbj BAC00817.1 phosphate transport system perm		8.0
gi 40255174 ref NP 775795.2 chromosome 10 open reading fra	<u>33</u> <u>33</u>	8.0
<u>gi 47213146 emb CAF93836.1 </u> unnamed protein product [Tetrao <u>gi 34905662 ref NP_914178.1 </u> P0475H04.14 [Oryza sativa (jap	<u>33</u> <u>33</u>	8.0 8.0

Alignments

Get selected sequences Select all Deselect all

```
gi|37182173|gb|AAQ88889.1| GRVS638 [Homo sapiens]
         Length = 140
 Score = 280 \text{ bits } (716), \text{ Expect} = 2e-74
 Identities = 140/140 (100%), Positives = 140/140 (100%)
 Frame = +2
Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT 268
          MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT
          MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT 60
Sbjct: 1
Query: 269 LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSAL 448
          LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSAL
Sbjct: 61 LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSAL 120
Ouery: 449 PAVTEMALFVTVFGLKKKPF 508
          PAVTEMALFVTVFGLKKKPF
Sbjct: 121 PAVTEMALFVTVFGLKKKPF 140
                                hypothetical protein MGC10744 isoform 1 [Homo sa
>gi|34101276|ref|NP 115730.2|
         Length = 146
 Score = 272 bits (695), Expect = 6e-72
 Identities = 139/146 (95%), Positives = 140/146 (95%), Gaps = 6/146 (4%)
 Frame = +2
Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDI-----QLV 250
          MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDI
Sbjct: 1
          MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIHPLPLCRLV 60
Query: 251 AALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIF 430
          AALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIF
Sbjct: 61 AALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIF 120
Query: 431 VFCSALPAVTEMALFVTVFGLKKKPF 508
          VFCSALPAVTEMALFVTVFGLKKKPF
Sbjct: 121 VFCSALPAVTEMALFVTVFGLKKKPF 146
gi|12834053|dbj|BAB22768.1| unnamed protein product [Mus musculus]
         Length = 140
 Score = 257 bits (657), Expect = 2e-67
 Identities = 125/140 (89%), Positives = 132/140 (94%)
 Frame = +2
Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT 268
          MGR+SGLVPSRFLTLLAHLVVVITLFWSR+SNIQACLPL FTPEEY+KQD QLVAAL +T
Sbjct: 1
          MGRISGLVPSRFLTLLAHLVVVITLFWSRESNIQACLPLKFTPEEYEKQDNQLVAALCLT 60
Query: 269 LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSAL 448
          LGLFAVELAGFLSGVSMFNSTQSL+SI AHCSASVALSFF+FERWECTTYWYIF FCSA
Sbjct: 61 LGLFAVELAGFLSGVSMFNSTQSLLSIAAHCSASVALSFFVFERWECTTYWYIFTFCSAF 120
Query: 449 PAVTEMALFVTVFGLKKKPF 508
```

PAVTE ALF+ VFGLKKKPF Sbjct: 121 PAVTETALFIAVFGLKKKPF 140

similar to RIKEN cDNA 1110004B13 [Rattus norvegi ->gi|27672842|ref|XP 213332.1| Length = 140Score = 257 bits (657), Expect = 2e-67Identities = 125/140 (89%), Positives = 132/140 (94%) Frame = +2Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT 268 MGR+SGLVPSRFLTLLAHLV+VITLFWSR+SNIQACLPL FTPEEY+KQD QLVAAL +T Sbjct: 1 MGRISGLVPSRFLTLLAHLVIVITLFWSRESNIQACLPLKFTPEEYEKQDNQLVAALCLT 60 Query: 269 LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSAL 448 LGLFAVELAGFLSGVSMFNSTQSL+SI AHCSASVALSFFIFERWECTTYWYIF FCSA Sbjct: 61 LGLFAVELAGFLSGVSMFNSTQSLLSIAAHCSASVALSFFIFERWECTTYWYIFAFCSAF 120 Query: 449 PAVTEMALFVTVFGLKKKPF 508 PAVTE ALF+ VFGLKKKPF Sbjct: 121 PAVTETALFIAVFGLKKKPF 140 □ >gi|47123437|gb|AAH70231.1| Unknown (protein for MGC:88213) [Homo sapiens] Length = 139Score = 245 bits (625), Expect = 8e-64Identities = 131/146 (89%), Positives = 132/146 (90%), Gaps = 6/146 (4%) Frame = +2Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDI-----QLV 250 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDI Sbjct: 1 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIHPLPLCRLV 60 Query: 251 AALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIF 430 AALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTY Sbjct: 61 AALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTY---- 116 Query: 431 VFCSALPAVTEMALFVTVFGLKKKPF 508 ALPAVTEMALFVTVFGLKKKPF Sbjct: 117 ---CALPAVTEMALFVTVFGLKKKPF 139 gi|12850002|dbj|BAB28560.1| unnamed protein product [Mus musculus] Length = 143Score = 217 bits (552), Expect = 2e-55Identities = 105/117 (89%), Positives = 111/117 (94%) Frame = +2Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT 268 MGR+SGLVPSRFLTLLAHLVVVITLFWSR+SNIQACLPL FTPEEY+KQD QLVAAL +T

MGRISGLVPSRFLTLLAHLVVVITLFWSRESNIQACLPLKFTPEEYEKQDNQLVAALCLT 60

Sbjct: 1

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Sbjct: 61 LGLFAVELAGFLSRVSMFNSTQSLLSIAAHCSASVALSFFVFERWECTTYWYIFTFC 117
qi|41351480|qb|AAH65897.1| Zqc:77926 [Danio rerio]
         Length = 135
 Score = 172 \text{ bits } (435), Expect = 8e-42
 Identities = 80/131 (61%), Positives = 100/131 (76%)
 Frame = +2
Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT 268
          M + LVP+RFLTL AHLV++IT+FWSRD+NIQ+CLPL FT ++Y +D +L ALSVT
          MSALKSLVPARFLTLTAHLVIIITIFWSRDNNIQSCLPLEFTEDQYRTEDTRLTVALSVT 60
Sbjct: 1
Query: 269 LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSAL 448
          L LF +ELAGFLSGVSMFNS Q+L+S+ H SA V LSFF+F +W C TYW IF CS
Sbjct: 61 LALFVLELAGFLSGVSMFNSNQALLSLITHSSACVCLSFFVFHQWPCWTYWIIFSICSVF 120
Query: 449 PAVTEMALFVT 481
          PAV E+ L ++
Sbjct: 121 PAVVELFLLLS 131
>gi|47225380|emb|CAG11863.1| unnamed protein product [Tetraodon nigroviridis]
         Length = 180
 Score = 169 \text{ bits } (429), \text{ Expect} = 4e-41
 Identities = 77/120 (64%), Positives = 100/120 (83%)
 Frame = +2
Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVT 268
          M VS LVP+RFLT++AHLV+VIT+FWSR++N++A LPL FT E+YD +D +LV AL+VT
Sbjct: 1
          MPAVSSLVPARFLTIIAHLVIVITIFWSRENNVRAGLPLDFTQEQYDSEDRKLVIALAVT 60
Query: 269 LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSAL 448
          +G+FA+ELAGF SGVSMFN +Q L+S G H SASVAL FF+FE+WEC YW+IFV C ++
Sbjct: 61 IGMFAIELAGFFSGVSMFNCSQGLLSTGTHASASVALLFFLFEQWECDIYWWIFVICRSV 120
Length = 141
 Score = 119 \text{ bits } (297), \text{ Expect = } 9e-26
 Identities = 70/113 (61%), Positives = 74/113 (65%), Gaps = 6/113 (5%)
 Frame = +2
Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDI-----QLV 250
          MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDI
Sbjct: 1
          MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIHPLPLCRLV 60
Query: 251 AALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWEC 409
          AALSVTLG
                           SG F
                                       + + H AS L
Sbjct: 61 AALSVTLG-----PLCSGAGRFPLRS--LHVQQHPRASSPLGLTVVHPWPC 104
```

Query: 269 LGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFC 439

LGLFAVELAGFLS VSMFNSTQSL+SI AHCSASVALSFF+FERWECTTYWYIF FC

```
Score = 114 \text{ bits } (284), \text{ Expect}(2) = 6e-33
Identities = 54/54 (100%), Positives = 54/54 (100%)
 Frame = +3
Query: 330 PRASSPLGLTVVHPWPCPSSYSSVGSALRIGTFLSSAVPFQLSLKWLYSSPSLG 491
          PRASSPLGLTVVHPWPCPSSYSSVGSALRIGTFLSSAVPFQLSLKWLYSSPSLG
Sbjct: 88 PRASSPLGLTVVHPWPCPSSYSSVGSALRIGTFLSSAVPFQLSLKWLYSSPSLG 141
Score = 50.1 bits (118), Expect(2) = 6e-33
 Identities = 21/21 (100%), Positives = 21/21 (100%)
 Frame = +1
Query: 271 GPLCSGAGRFPLRSLHVQQHP 333
          GPLCSGAGRFPLRSLHVQQHP
Sbjct: 68 GPLCSGAGRFPLRSLHVQQHP 88
- >gi|34535000|dbj|BAC87177.1| unnamed protein product [Homo sapiens]
         Length = 163
 Score = 79.0 \text{ bits (193)}, Expect = 1e-13
 Identities = 33/33 (100%), Positives = 33/33 (100%)
 Frame = +3
Query: 246 WWPRSLSPWASLQWSWPVSSQESPCSTAPRASS 344
          WWPRSLSPWASLQWSWPVSSQESPCSTAPRASS
Sbjct: 84 WWPRSLSPWASLQWSWPVSSQESPCSTAPRASS 116
 Score = 60.8 bits (146), Expect = 3e-08
 Identities = 32/41 (78%), Positives = 32/41 (78%)
 Frame = \pm 2
Query: 89 MGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTF 211
           MGRVSGLVPSRFLTLLAHLVVVITLFWSR
         MGRVSGLVPSRFLTLLAHLVVVITLFWSRVRPTAALNPSPF 41
Sbjct: 1
□>gi|7500843|pir||T21990 hypothetical protein F39B2.9 - Caenorhabditis elegans
         Length = 136
 Score = 37.7 bits (86), Expect = 0.25
 Identities = 31/127 (24%), Positives = 58/127 (45%)
 Frame = +2
Query: 122 FLTLLAHLVVVITLFWSRDSNIQACLPLTFTPEEYDKQDIQLVAALSVTLGLFAVELAGF 301
                          +S+S+++A++E+I
                                                        L T
                                                                 +E A
           F L+ H +
Sbjct: 12 FQCLMGHFALSTITIFSQASHLEASVSGGASISEEAR--IGFTVCLVFTKLAILIEFASI 69
Query: 302 LSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFVFCSALPAVTEMALFVT 481
          L V + + ++SL S +H AS+
                                      FI++
                                                 +W +F F S P V +A+ +
```

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Sbjct: 70 L--VDLPSPSRSLYSTLSHTIASIFFLIFIYDSHPPAHFWLLFAFLSLPPCV--LAVSAS 125
Query: 482 VFGLKKK 502
          + G ++K
Sbjct: 126 IGGFRRK 132
>gi|46362710|ref|ZP 00225559.1| hypothetical protein Krad06004793 [Kineococcus r
          SRS30216]
         Length = 293
 Score = 37.4 \text{ bits } (85), \text{ Expect = } 0.33
 Identities = 31/87 (35%), Positives = 41/87 (47%)
 Frame = +1
Query: 52 PASETAALVPEGHGPGLRACALSLPDAPGASGGRHHLILVPGQQHTGLPASHVHPRGV*Q 231
          P + A+ PEG G GL A+ L + PGA G +H +L GQ+
Sbict: 57 PVPDGVAVHPEGVGGGLPG-AVVLEEGPGA-GPQHGPVLDRGQRRQPPPRPRPHRAGVQD 114
Query: 232 AGHSAGGRALCHPGPLCSGAGRFPLRS 312
                      PGP
                              G+ P+R+
           G AG
Sbjct: 115 LGQRAG-----PGP-----GQRPVRA 130
- >gi|18025542|gb|AAF78882.2| EBNA-3B [cercopithicine herpesvirus 15]
         Length = 928
 Score = 36.6 \text{ bits } (83), \text{ Expect = } 0.56
 Identities = 42/149 (28%), Positives = 50/149 (33%), Gaps = 17/149 (11%)
 Frame = -2
Query: 407 TPNARI*RRTGPR----- 258
                              + + P W
           TP A I
                    + PR
                                                              PG R
Sbjct: 601 TPRAHIPANSDPRRATFDVLGFRSPDWPPKNWTNANPSTANANASRPISAAPGRARPPAA 660
Query: 257 ----ARPPAECPACHTPRG*T*EAGRPVCCCPGTRIR**RPPDAPGASGSERAQALRPG- 93
                                    RP
                                          PG
                                                  RPP A A G R A PG
               ARPPA P
Sbjct: 661 APGRARPPAAAPGRARPPAAATGRARPPAAAPGRA----RPPAA--APGRARPPAAAPGR 714
          --PWPSGTRAAVSEAGSSETATPEVSPQA 12
Query: 92
                         A +
                               A P+ PQA
                     A
Sbjct: 715 ARPPAAAPGRARPPAAAPGRAAPQQQPQA 743
                                 similar to Hypothetical protein MGC37938 [Homo s
->gi|41114515|ref|XP 371222.1|
         Length = 215
 Score = 36.2 \text{ bits } (82), \text{ Expect = } 0.73
 Identities = 34/110 (30%), Positives = 45/110 (40%), Gaps = 1/110 (0%)
 Frame = +1
Query: 28 SGVAVSELPASETAALVPEGHG-PGLRACALSLPDAPGASGGRHHLILVPGQQHTGLPAS 204
                       + L+ G G PG R A +LP P +G R L
           + + V E
Sbjct: 92 AALVVLEFSLRAVSTLLSLGKGLPGCRREAAALPAVPVLAGLRADLW--PELPAGGRPSP 149
Query: 205 HVHPRGV*QAGHSAGGRALCHPGPLCSGAGRFPLRSLHVQQHPEPHLHWG 354
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+AGH+AG P P PLR+ +P L WG
          H P
Sbjct: 150 HAEPAAEPRAGHAAGPGCPAPPPPRLP----PLRAAQ-----QPALLWG 189
\square >gi|46321304|ref|ZP 00221682.1| COG0477: Permeases of the major facilitator supe
           [Burkholderia cepacia R1808]
         Length = 428
Score = 35.8 bits (81), Expect = 0.95
Identities = 22/77 (28%), Positives = 38/77 (49%), Gaps = 1/77 (1%)
Frame = +2
Query: 254 ALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFV 433
          A+SV + L+ + + + F ++++
                                              SA VALS + + W+
Sbjct: 57 AISVNIALYGLTGPFAAAAMQRFGLRPTILTALVTLSAGVALSSMMTQSWQMVVVWGLMV 116
Query: 434 FCS-ALPAVTEMALFVT 481
           CS + A+T A FVT
Sbjct: 117 GCSTGVVALTLSATFVT 133
- >gi|47210644|emb|CAG06315.1| unnamed protein product [Tetraodon nigroviridis]
         Length = 325
Score = 35.4 bits (80), Expect = 1.2
Identities = 28/88 (31%), Positives = 41/88 (46%)
Frame = -3
Query: 280 KEAQGDRERGHQLNVLLVILLGGERERQAGLYVAVPGPE*GDDDHQMRQERQEARGHKP* 101
          + A+G R + H+L+ + + GG + GL
                                            VPGP
                                                     DH++R+ER
                                                                +G
Sbjct: 203 RRARGRRGQEHRLHAGVPLGQGGLQ----GLRRPVPGPVRALRDHEVREERARGQGRAVR 258
Query: 100 DPAHGPRGQGRRSLRLEVQRQRLLKSPR 17
                            QR + L PR
            A P G
                     R
Sbjct: 259 AGAEPPAGGAHRR-----QRAQRLHPPR 281
                                  unknown protein [Oryza sativa (japonica cultivar-g
__>gi|34902106|ref|NP 912399.1|
gi|29893617|gb|AAP06871.1|
                             unknown protein [Oryza sativa (japonica cultivar-group
         Length = 279
 Score = 35.4 bits (80), Expect = 1.2
Identities = 18/46 (39%), Positives = 25/46 (54%)
Frame = +3
Query: 267 PWASLQWSWPVSSQESPCSTAPRASSPLGLTVVHPWPCPSSYSSVG 404
          PW SL+ S PV +E P + AP + + GLT+
Sbjct: 164 PWLSLRNSCPVCRRELPAAAAPESEADAGLTI---WRLPRGGFAVG 206
__>gi|7106228|gb|AAF36091.1|
                               flagelliform silk protein [Nephila madagascariensis]
         Length = 1884
Score = 35.0 \text{ bits } (79), \text{ Expect = } 1.6
Identities = 28/96 (29%), Positives = 35/96 (36%), Gaps = 2/96 (2%)
```

Frame = +1Query: 10 SACGETSGVAVSELPASETAALVPEGHGPGLRACALSLPDAPGASGGRHHLILVPGQQHT 189 P G GPG + S P S G+ SG G SG R Sbjct: 118 SGAGQPSGAGPSGTGGYAPTGYAPSGSGPGGVRPSASGPSGSGPSGSRPSSSGSSGTRPS 177 Query: 190 GLPASHVHPRGV*QAGHSAGGRAL--CHPGPLCSGA 291 GSGG+ P G+ GP SG+ Sbjct: 178 ANAAGGSSPGGIAPGGSSPGGAGVSGATGGPASSGS 213 $\square > gi | 34536289 | dbj | BAC87601.1 |$ unnamed protein product [Homo sapiens] Length = 181Score = 34.7 bits (78), Expect = 2.1Identities = 26/76 (34%), Positives = 30/76 (39%), Gaps = 14/76 (18%) Frame = +3Query: 249 WPRSLS-----PWASLQWSWPVSSQESPCSTAPRASSPLGLT---VVHPWPCPS---- 386 P + L W WPWP S+ P ASLL V PWPCP+ Sbjct: 102 WPCSVFGRALCYPSPALPWPWPYHGLLLPWPGPTLAFSTLALPFPGVALPWPCPALALAL 161 Query: 387 SYSSVGSALRIGTFLS 434 GSAL + LS Sbjct: 162 PYPGPGSALTLALVLS 177 - >gi|6686017|sp|062732|SYN1 CANFA Synapsin I gi|2944066|gb|AAC05207.1| synapsin I [Canis familiaris] Length = 415Score = 34.7 bits (78), Expect = 2.1Identities = 31/99 (31%), Positives = 43/99 (43%), Gaps = 2/99 (2%) Frame = -2Query: 305 RGNRPAPLQRGPG*QRARPPAECPACHTPRG*T*EAGRPVCCCPGTRIR**RPPDAPGA- 129 +G + P+ GPG APAPA+P+ +AG P PP A GA T +Sbjct: 257 QGRQSRPVAGGPG---APPAARPPASPSPQR---QAGPPQATRQ-TSVSGQAPPKASGAP 309 Query: 128 -SGSERAQALRPGPWPSGTRAAVSEAGSSETATPEVSPQ 15 SG +R + P P+G S+AG Sbjct: 310 PSGQQRQGPPQKPPGPAGPTRQASQAGPMPRTGPPTTQQ 348 \square >gi|14701574|dbj|BAB62028.1| elastase precursor [Prevotella intermedia] Length = 640Score = 34.7 bits (78), Expect = 2.1Identities = 18/59 (30%), Positives = 25/59 (42%) Frame = +3Query: 249 WPRSLSPWASLQWSWPVSSQESPCSTAPRASSPLGLTVVHPWPCPSSYSSVGSALRIGT 425 WP + PWWP + P + P+P P Y +G +R GTP+G Sbjct: 265 WPWPVDPW-----WPPIPKPWP----PIGPGPIGPGPIGPYPPPRPYRPIGPRIRFGT 313

```
Length = 2064
 Score = 34.3 \text{ bits } (77), \text{ Expect} = 2.8
 Identities = 19/44 (43%), Positives = 27/44 (61%), Gaps = 2/44 (4%)
 Frame = +3
Query: 258 SLSPWASLQWSWPVSSQESPC--STAPRASSPLGLTVVHPWPCP 383
           SLSP S + S P +SQ+ PC S+AP + +P+G + WP P
Sbjct: 2006 SLSPDVSTEASPPRASQDIPCLDSSAPESGTPMG--ALGDWPAP 2047
                              hypothetical protein [Homo sapiens]
_>gi|34365329|emb|CAE45990.1|
         Length = 924
 Score = 34.3 bits (77), Expect = 2.8
 Identities = 19/44 (43%), Positives = 27/44 (61%), Gaps = 2/44 (4%)
 Frame = +3
Query: 258 SLSPWASLQWSWPVSSQESPC--STAPRASSPLGLTVVHPWPCP 383
          SLSP S + S P +SQ+ PC S+AP + +P+G
Sbjct: 866 SLSPDVSTEASPPRASQDIPCLDSSAPESGTPMG--ALGDWPAP 907
\square >gi|46311822|ref|ZP 00212424.1| COG0477: Permeases of the major facilitator supe
          [Burkholderia cepacia R18194]
         Length = 427
 Score = 34.3 bits (77), Expect = 2.8
 Identities = 21/77 (27%), Positives = 38/77 (49%), Gaps = 1/77 (1%)
 Frame = +2
Query: 254 ALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFV 433
          A+SV + L+ + + F
                                   ++++
                                             A VALS + + W+
                                                              W + V
Sbjct: 57 AISVNIALYGLTGPFAAAAMQRFGLRPTILTALVTMCAGVALSSMMTQSWQMVVIWGLMV 116
Query: 434 FCSA-LPAVTEMALFVT 481
           CS+ + A+T A FVT
Sbjct: 117 GCSSGVVALTLSATFVT 133
- >gi|13562004|gb|AAK30605.1| major ampullate spidroin 2-like protein [Nephila mad
         Length = 1953
 Score = 34.3 bits (77), Expect = 2.8
 Identities = 26/85 (30%), Positives = 36/85 (42%), Gaps = 1/85 (1%)
 Frame = +1
Query: 52 PASETAALVPEGHGPGLRACALSLPDAPGASGGRHHLILVPGQQHTGLPASHVHPRGV*Q 231
           P + AA P G+GPG + A G+ G +
                                               PGQQ G P +
Sbjct: 1562 PGAAAAAAGPGGYGPGQQGPGAAAAAAAGSGPGGYG----PGQQGPGGPGAAAAAAGRGP 1617
Query: 232 AGHSAGGRALCHPG-PLCSGAGRFP 303
            G+ G + PG
                            + AGR P
Sbjct: 1618 GGYGPGQQGPGGPGAAAAAAAGRGP 1642
```

```
Score = 34.3 bits (77), Expect = 2.8
 Identities = 26/85 (30%), Positives = 36/85 (42%), Gaps = 1/85 (1%)
 Frame = +1
           PASETAALVPEGHGPGLRACALSLPDAPGASGGRHHLILVPGQQHTGLPASHVHPRGV*Q 231
Query: 52
            P + AA P G+GPG + + A G+ G +
                                                    PGQQ G P +
Sbjct: 951 PGAAAAAAGPGGYGPGQQGPGAAAAAAAGSGPGGYG----PGQQGPGGPGAAAAAAGRGP 1006
Query: 232 AGHSAGGRALCHPG-PLCSGAGRFP 303
                     PG + AGR P
             G+ G+
Sbjct: 1007 GGYGPGQQGPGGPGAAAAAAAAGRGP 1031
- >gi|28279307|gb|AAH46165.1| Unknown (protein for IMAGE:5192014) [Homo sapiens]
 gi|29387251|gb|AAH48278.1| Unknown (protein for IMAGE:5192302) [Homo sapiens]
          Length = 205
 Score = 33.9 \text{ bits } (76), \text{ Expect = } 3.6
 Identities = 20/40 (50%), Positives = 21/40 (52%), Gaps = 5/40 (12%)
 Frame = -2
Query: 134 GASGSERAQAL-RPGPWPSGTRAAVSEAGS----SETATP 30
                                     +A S
           GA G ER L RPG WP G RA
                                             S T TP
Sbjct: 59 GAGGGEREDGLWRPGRWPRGARAPGEQAASRSPWSATGTP 98
- >qi|34495966|ref|NP 900181.1| probable MFS permease [Chromobacterium violaceum A
 qi|34101820|qb|AAQ58188.1| probable MFS permease [Chromobacterium violaceum ATCC
          Length = 425
 Score = 33.9 \text{ bits } (76), \text{ Expect = } 3.6
 Identities = 23/77 (29%), Positives = 36/77 (46%)
 Frame = +2
Query: 254 ALSVTLGLFAVELAGFLSGVSMFNSTQSLISIGAHCSASVALSFFIFERWECTTYWYIFV 433
           ALS+ L LF +
                            + + F
                                     ++++S A
                                                ASVALS +
                                                                   W + V
Sbjct: 57 ALSINLALFGLMGPFAAAAMLRFGLRRTVLSALALLGASVALSTLMRSNWQLQLLWGVMV 116
Query: 434 FCSALPAVTEMALFVTV 484
                   T M L + V
            C+
Sbjct: 117 GCAT--GATAMTLGASV 131
>gi|37680831|ref|NP 935440.1| hypothetical protein VV2647 [Vibrio vulnificus YJ0
                               hypothetical protein [Vibrio vulnificus YJ016]
 gi|37199580|dbj|BAC95411.1|
          Length = 464
 Score = 33.9 \text{ bits } (76), \text{ Expect} = 3.6
 Identities = 29/75 (38%), Positives = 39/75 (52%), Gaps = 3/75 (4%)
 Frame = +2
Query: 74 LSPRAMGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFT-PEEYDKQDIQLV 250
                   VSG++ S
                              L+A VV
                                        LFWSR S
                                                  A LPL
                                                                    + L+
           LS A
```

```
Sbjct: 361 LSGIAYSLVSGVLDSPLSQLMA--VVACALFWSRCSLASASLPLPANKPRRWWAHGVILI 418
Query: 251 AAL--SVTLGLFAVE 289
           A+ SV++G A+E
Sbjct: 419 IAIGCSVSVGYRALE 433
>gi|27365118|ref|NP 760646.1| Unknown [Vibrio vulnificus CMCP6]
 gi|27361264|gb|AA010173.1| Unknown [Vibrio vulnificus CMCP6]
         Length = 464
 Score = 33.9 \text{ bits } (76), \text{ Expect = } 3.6
 Identities = 29/75 (38%), Positives = 39/75 (52%), Gaps = 3/75 (4%)
 Frame = +2
Query: 74 LSPRAMGRVSGLVPSRFLTLLAHLVVVITLFWSRDSNIQACLPLTFT-PEEYDKQDIQLV 250
          LS A VSG++ S L+A VV LFWSR S A LPL
                                                           P +
Sbjct: 361 LSGIAYSLVSGVLDSPLSQLMA--VVACALFWSRCSLASASLPLPANKPRRWWAHGVILI 418
Query: 251 AAL--SVTLGLFAVE 289
           A+ SV++G A+E
Sbjct: 419 IAIGCSVSVGYRALE 433
- >gi|24575123|gb|AAL06694.1| putative regulatory protein [Streptomyces globisporu
         Length = 196
 Score = 33.9 \text{ bits } (76), \text{ Expect = } 3.6
 Identities = 23/52 (44%), Positives = 26/52 (50%), Gaps = 6/52 (11%)
 Frame = +1
Query: 64 TAALVPEGHGPGLRACALSLPDAPGA----SGGRHHLILVPGQQHTGLPA 201
                                           S GR
           TA + +GH PGL A LS+PDA G
Sbjct: 146 TAQAMEDGH-PGLEAMGLSMPDADGLWRFKDYSYGREFARLHPDTTSTGSPA 196
>gi|41057143|ref|NP 957857.1| ORF080 virion core protein [Orf virus]
 gi|41018700|gb|AAR98305.1| ORF080 virion core protein [Orf virus]
         Length = 328
 Score = 33.9 \text{ bits } (76), \text{ Expect} = 3.6
 Identities = 28/94 (29%), Positives = 34/94 (36%), Gaps = 4/94 (4%)
 Frame = -2
Query: 293 PAPLQRGPG*QRARPPAECPACHTPRG*T*EAGRPVCCCPGTRIR**RPP----DAPGAS 126
                      A PPA P C P
                                               C P TR
                                         Α
Sbjct: 155 PAPAVTCPAPAPACPPATAPTCPPPAVCPAPARPAPACPPSTRQCPPAPPLPTKPAPAAK 214
Query: 125 GSERAQALRPGPWPSGTRAAVSEAGSSETATPEV 24
               Q+L P +P+ + +S
Sbjct: 215 PIFLQQSLPPPQYPASSCPTIKAPAASPVLEPRV 248
☐ >gi|47230172|emb|CAG10586.1|
                                 unnamed protein product [Tetraodon nigroviridis]
         Length = 441
```

```
Score = 33.5 \text{ bits } (75), \text{ Expect = } 4.7
 Identities = 40/118 (33%), Positives = 48/118 (40%), Gaps = 13/118 (11%)
 Frame = +1
Query: 7 LSACGETSGVAVSELPASETAALVPEGH--GPGLRACALSLPDAPGA--SGGRHHLILVP 174
           L A GE
                                      H GP RA
                                                    P+APG+ +G +
                         +P
Sbjct: 256 LPAGGEREQRAAGRVPGGPLRGHPAVAHQRGPLPRAH----PEAPGSPQAGDQRGAAPVP 311
Query: 175 -GQQHTGLPASHVHPRGV*QAGHSAGGRALCHP-----GPLCSGAGRFPLRSLHV 321
            G QH GLP PR + AG +A G
                                        P PL +GA R P R HV
Sbjct: 312 RGAQHAGLPQHAAQPRPLLPAGQAALGLPGLRPRARLSQLGQPLGAGAQR-PARVPHV 368
- >qi|39583515|emb|CAE73973.1| Hypothetical protein CBG21601 [Caenorhabditis brigg
         Length = 719
 Score = 33.5 \text{ bits } (75), \text{ Expect = } 4.7
 Identities = 24/90 (26%), Positives = 30/90 (33%)
 Frame = -2
Query: 290 APLQRGPG*QRARPPAECPACHTPRG*T*EAGRPVCCCPGTRIR**RPPDAPGASGSERA 111
           A \quad R \quad P \quad + \quad + \quad P \quad A \quad P
                                  PR
                                                 PG
                                                             AP AS
Sbjct: 548 ASAPRAPAPRASAPRASAPRASAPRASAPRASAPRALAPGA-----SAPSASAPRAS 599
Query: 110 QALRPGPWPSGTRAAVSEAGSSETATPEVS 21
                 P S TRA+ A + P S
Sbjct: 600 APRASAPRASAPRASAPRASAPRAS 629
>qi|9663052|emb|CAC01096.1| 110-R orphan receptor [Haemonchus contortus]
          Length = 986
 Score = 33.1 \text{ bits } (74), \text{ Expect = } 6.1
 Identities = 16/44 (36%), Positives = 23/44 (52%)
 Frame = -2
Query: 149 PPDAPGASGSERAQALRPGPWPSGTRAAVSEAGSSETATPEVSP 18
            PP P A G+ ++ +RP +A S+ GSS
                                                  T EV+P
Sbjct: 921 PPPPPPAQGTTPSKVIRPPSSKMSDDSAYSDGGSSSVLTTEVTP 964
[] >gi|47214200|emb|CAG00828.1| unnamed protein product [Tetraodon nigroviridis]
         Length = 2135
 Score = 33.1 \text{ bits } (74), \text{ Expect = } 6.1
 Identities = 34/105 (32%), Positives = 42/105 (40%), Gaps = 5/105 (4%)
 Frame = +1
Query: 1 LDLSACGETSGVAVSELPASETAALVPEGH--GPGLRACALSLPDAPGASGGRHHLILVP 174
           LL+G+SG+ LP PE H GP L LS P PG S
Sbjct: 1484 LGLPSAGASSGSSAFGLPP-----PELHLRGPSLSQNKLSRPHEPGLSDSPSSLPNVP 1536
Query: 175 GQ-QHTGLPASHVHPRGV*QAGHSAGGRALCHP--GPLCSGAGRF 300
                H + +SH + PG + H
                                        A HP P G+ F
Sbjct: 1537 SPGTHFHISSSHLQPSG---SSHPQLSLAGLHPLLPPAGGGSAEF 1578
```

```
>gi|47217474|emb|CAG10243.1| unnamed protein product [Tetraodon nigroviridis]
          Length = 1190
 Score = 33.1 bits (74), Expect = 6.1
 Identities = 22/62 (35%), Positives = 28/62 (45%), Gaps = 4/62 (6%)
 Frame = -2
Query: 194 RPVCCCPGTRIR**RPPDAPGA----SGSERAQALRPGPWPSGTRAAVSEAGSSETATPE 27
                                                GP +G
                         RPP APGA
                                      GS A
                                                             + GS+E+ P
                CP R
           RP
Sbjct: 97 RPRASCPRLR----RPPGAPGAHCSRQGSAEASPSWAGPSHAGFLPPTRKPGSNESIPPA 152
Query: 26 VS 21
           ++
Sbjct: 153 IT 154
>gi|46432590|gb|EAK92065.1| hypothetical protein CaO19.6598 [Candida albicans SC
          Length = 781
 Score = 33.1 \text{ bits } (74), \text{ Expect = } 6.1
 Identities = 28/85 (32%), Positives = 34/85 (40%), Gaps = 3/85 (3%)
 Frame = -2
Query: 251 PPAEC-PACHTPRG*T*EAGRPVCCCPGT--RIR**RPPDAPGASGSERAQALRPGPWPS 81
                      PR T + G P
                                     PG
                                               +PP AP
                                                                  PPP
           PPA
Sbjct: 475 PPARATPPAPPPRARTNQLGLPPRNTPGLPPRTNNTQPPPAPPPRASRGAVPPPPPPPPR 534
Query: 80 GTRAAVSEAGSSETATPEVSPQADK 6
            TRA +
                      S + +SP A +
Sbjct: 535 ATRAPMQLQLQSSPQSSPISPPAQQ 559
- >qi|15220207|ref|NP 172545.1|
                                  expressed protein [Arabidopsis thaliana]
 gi|28393599|gb|AA042219.1|
                              putative carboxyl-terminal peptidase [Arabidopsis thal
 gi|29824323|gb|AAP04122.1|
                              putative carboxyl-terminal peptidase [Arabidopsis thal
          Length = 467
 Score = 32.7 bits (73), Expect = 8.0
 Identities = 15/31 (48%), Positives = 16/31 (51%)
Frame = -3
Query: 148 HQMRQERQEARGHKP*DPAHGPRGQGRRSLR 56
               +
                     RG KP DP
                                PRG RR LR
           НО
Sbjct: 135 HQPAFDHPSLRGQKPLDPPERPRGHNRRGLR 165
☐ >gi|46443901|qb|EAL03180.1|
                                hypothetical protein CaO19.4050 [Candida albicans SC
gi|46444066|gb|EAL03344.1|
                              hypothetical protein CaO19.11532 [Candida albicans SC5
         Length = 154
Score = 32.7 \text{ bits } (73), \text{ Expect} = 8.0
Identities = 16/49 (32%), Positives = 25/49 (51%)
 Frame = +3
```

```
Query: 255 RSLSPWASLQWSWPVSSQESPCSTAPRASSPLGLTVVHPWPCPSSYSSV 401
                      +W +S S + +P SSP
                                              T++ WPC
           R +S +
Sbjct: 2
           REMSSMSESSLAWTAASSSSLVTNSPSFSSPWPNTLIRNWPC--GYSSL 48
>qi|21388708|dbj|BAC00817.1| phosphate transport system permease ABC transporter
           [Rhodobacter sphaeroides f. sp. denitrificans]
          Length = 481
 Score = 32.7 \text{ bits } (73), \text{ Expect} = 8.0
 Identities = 31/102 (30%), Positives = 40/102 (39%)
 Frame = +1
Query: 34 VAVSELPASETAALVPEGHGPGLRACALSLPDAPGASGGRHHLILVPGQQHTGLPASHVH 213
                    + AAL PEG P LRA
                                       ++ D
                                                            + H L AS
Sbjct: 105 VVAEGLSDRDLAALTPEGLAPRLRAVGAAVADITPQT-----LDAARAHRALQASFGL 157
Query: 214 PRGV*QAGHSAGGRALCHPGPLCSGAGRFPLRSLHVQQHPEP 339
                  G S GGR+
                             PGP
                                     AG
                                           + +
                                                  H + P
Sbjct: 158 WMSVLVLGASLGGRS---PGPPAHSAGLPCPQRVRALHHVDP 196
- >gi|40255174|ref|NP 775795.2| Chromosome 10 open reading frame 64 [Homo sapien
                              unnamed protein product [Homo sapiens]
gi|34534655|dbj|BAC87073.1|
          Length = 1042
 Score = 32.7 \text{ bits } (73), \text{ Expect} = 8.0
 Identities = 13/25 (52%), Positives = 16/25 (64%)
 Frame = -2
Query: 137 PGASGSERAQALRPGPWPSGTRAAV 63
            PG SGS+ AQ L GPWP+
Sbjct: 951 PGCSGSQTAQGLAEGPWPAAPDAGL 975
□ >gi|47213146|emb|CAF93836.1| unnamed protein product [Tetraodon nigroviridis]
         Length = 708
Score = 32.7 \text{ bits } (73), \text{ Expect} = 8.0
Identities = 18/38 (47%), Positives = 20/38 (52%), Gaps = 1/38 (2%)
Frame = +1
Query: 79 PEGHGPGLRACALSLPDAPGA-SGGRHHLILVPGQQHT 189
           PEG GP +A A S P+ P
                                  GGR L
                                           P Q HT
Sbjct: 508 PEGEGPRTKAGACSTPEGPATPPGGRKILKKTPEQLHT 545
-gi|34905662|ref|NP 914178.1| P0475H04.14 [Oryza sativa (japonica cultivar-group
gi|12060519|dbj|BAB20648.1| P0475H04.14 [Oryza sativa (japonica cultivar-group)]
         Length = 529
Score = 32.7 \text{ bits } (73), \text{ Expect} = 8.0
Identities = 22/66 (33%), Positives = 27/66 (40%), Gaps = 2/66 (3%)
```

Frame = +3

Query: 279 LQWSWPVSS--QESPCSTAPRASSPLGLTVVHPWPCPSSYSSVGSALRIGTFLSSAVPFQ 452

L WS+P S + PC P+ P PCP Y V AL + SA P

Sbjct: 68 LSWSFPPSIVLEHHPCENFSFFPPPIDRKRTGPRPCPVCYVPVEQALALMPGAPSASPVL 127

Query: 453 LSLKWL 470

SL +L

Sbjct: 128 RSLNYL 133

Get selected sequences Select all Deselect all

Lambda K H

0.318 0.134 0.401

Gapped

Lambda K H

0.267 0.0410 0.140

Matrix: BLOSUM62

Gap Penalties: Existence: 11, Extension: 1

Number of Sequences: 1866121

Number of Hits to DB: 192,225,388

Number of extensions: 5047647

Number of successful extensions: 16879

Number of sequences better than 10.0: 9

Number of HSP's better than 10.0 without gapping: 15304

Number of HSP's gapped: 16825

Number of HSP's successfully gapped: 10

Number of extra gapped extensions for HSPs above 10.0: 0

Length of query: 759

Length of database: 619,474,291

Length adjustment: 122

Effective length of query: 637

Effective length of database: 391,807,529

Effective search space: 249581395973

Effective search space used: 51326786299

T: 12

A: 40

X1: 16 (7.3 bits)

X2: 38 (15.0 bits)

X3: 64 (25.0 bits)

S1: 41 (22.0 bits)

S2: 72 (32.3 bits)